## Amendments to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

## Listing of Claims:

- 1-32 (canceled)
- 33. (currently amended) A method for isolating an miRNA a microRNA of interest from a sample comprising the miRNA microRNA of interest; the method comprising:
  - a) providing a sample comprising the miRNA microRNA of interest;
  - b) providing a capture probe according to claim 1 comprising:
    - i) a first adapter segment having a first adapter segment sequence, the first adapter segment comprising a 3' end and a 5' end;
    - ii) a second adapter segment having a second adapter segment sequence, the second adapter segment comprising a 3' end and a 5' end; and
    - iii) a microRNA binding segment having a microRNA binding segment sequence:
    - where the microRNA binding segment is substantially complementary to, and capable of hybridizing to, one or more than one microRNA of interest by Watson-Crick base pairing;
    - where the 5' end of the first adapter segment is connected to the 3' end of the microRNA binding segment; and
    - where the 3' end of the second adapter segment is connected to the 5' end of the microRNA binding segment;
  - c) providing a first linker and a second linker;
  - d) combining the sample, the capture probe, the first linker and the second linker;

e) allowing the first linker to hybridize with the first adapter segment, the miRNA microRNA of interest to hybridize with the miRNA microRNA binding segment, and the second linker to hybridize with the second adapter segment;

- f) ligating the 3' end of the first linker that is hybridized to the first adapter segment to the 5' end of the miRNA microRNA of interest that is hybridized to the miRNA microRNA binding segment, and ligating the 3' end of the miRNA microRNA of interest that is hybridized to the miRNA microRNA binding segment to the 5' end of the second linker that is hybridized to the second adapter segment, thereby producing a complex defined as a strand of first linker, miRNA microRNA of interest and second linker that have been ligated together (ligated first linker-miRNA microRNA of interest-second linker) and that is hybridized to the capture probe; and
- g) dehybridizing the capture probe from the strand of the ligated first linker-miRNA microRNA of interest-second linker;

where the miRNA microRNA of interest has an miRNA a microRNA of interest sequence, and comprises a 3' end and a 5' end;

where the miRNA microRNA of interest is substantially complementary to, and capable of hybridizing to, the miRNA microRNA binding segment of the capture probe by Watson-Crick base pairing;

where the first linker has a first linker sequence, and comprises a 3' end and a 5' end; where the first linker is substantially complementary to, and capable of hybridizing to, the first adapter segment of the capture probe by Watson-Crick base pairing;

where the second linker has a second linker sequence, and comprises a  $3^{\circ}$  end and a  $5^{\circ}$  end; and

where the second linker is substantially complementary to, and capable of hybridizing to, the second adapter segment of the capture probe by Watson-Crick base pairing.

34. (currently amended) The method of claim 33, where the sample further comprises one or more than one substance that is chemically related to the miRNA microRNA of interest selected from the group consisting of an RNA other than a miRNA microRNA and a DNA.

- 35. (original) The method of claim 33, where the sample is from a eukaryote.
- 36. (original) The method of claim 33, where the sample is from a primate.
- 37. (original) The method of claim 33, where the sample is from a human.
- 38. (original) The method of claim 33, where the sample comprises a tissue or fluid selected from the group consisting of blood, brain, heart, intestine, liver, lung, pancreas, muscle, a leaf, a flower, a plant root and a plant stem.
- 39. (currently amended) The method of claim 33, where the miRNA microRNA of interest consists of 18 or 19 or 20 or 21 or 22 or 23 or 24 RNA residues.
- 40. (currently amended) The method of claim 33, where the miRNA microRNA of interest is listed in a public database.
- 41. (currently amended) The method of claim 33, where the sample provided comprises a plurality of miRNAs microRNAs of interest; and

where each of the plurality of miRNAs microRNAs of interest has miRNA microRNA of interest sequences that are identical to one another.

- 42. (currently amended) The method of claim 33, where the sample provided comprises a plurality of miRNAs microRNAs of interest comprising a first miRNA microRNA of interest having a first miRNA microRNA of interest sequence, and a second miRNA microRNA of interest having a second miRNA microRNA of interest sequence; and
- where the first miRNA microRNA of interest sequence is different from the second miRNA microRNA of interest sequence.
- 43. (currently amended) The method of claim 33, where the sample provided comprises a plurality of miRNAs microRNAs of interest comprising a first miRNA microRNA of interest sequence, a second miRNA microRNA of interest sequence, and a third miRNA microRNA of interest sequence, and a third miRNA microRNA of interest sequence;

where the first miRNA microRNA of interest sequence is different from the second miRNA microRNA of interest sequence;

where the first miRNA microRNA of interest sequence is different from the third miRNA microRNA of interest sequence; and

where second miRNA microRNA of interest sequence is different from the third miRNA microRNA of interest sequence.

- 44. (original) The method of claim 33, further comprising isolating the total RNA from the sample after providing the sample.
- 45. (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes;

where each of the capture probes comprises identical first adapter segment sequences; where each of the capture probes of the set of capture probes comprises identical miRNA microRNA binding segment sequences; and

where each of the capture probes of the set of capture probes comprises identical second adapter segment sequences.

46. (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes;

where the set comprises at least one capture probe comprising an miRNA a microRNA binding segment that is substantially complementary to, and capable of hybridizing to, each miRNA microRNA listed in a single public database.

 (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes;

where the set comprises a first capture probe and a second capture probe;

where the first capture probe and the second capture probe have identical first adapter segment sequences;

where the first capture probe and the second capture probe have identical  $\frac{miRNA}{microRNA}$  binding segment sequences; and

where the first capture probe has a second adapter segment sequence that is different from the second adapter segment sequence of the second capture probe.

48. (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes;

where the set comprises a first capture probe and a second capture probe;

where the first capture probe and the second capture probe have identical first adapter segment sequences;

where the first capture probe and the second capture probe have identical second adapter segment sequences; and

where the first capture probe has an miRNA a microRNA binding segment sequence that is different from the miRNA microRNA binding segment sequence of the second capture probe.

49. (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes;

where the set comprises a first capture probe and a second capture probe;

where the first capture probe and the second capture probe have identical miRNA microRNA binding segment sequences;

where the first capture probe and the second capture probe have identical second adapter segment sequences; and

where the first capture probe has a first adapter segment sequence that is different from the first adapter segment sequence of the second capture probe.

50. (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes:

where the set comprises a first capture probe and a second capture probe;

where the first capture probe and the second capture probe have identical first adapter segment sequences;

where the first capture probe has an miRNA a microRNA binding segment sequence that is different from the miRNA microRNA binding segment sequence of the second capture probe; and

where the first capture probe has a second adapter segment sequence that is different from the second adapter segment sequence of the second capture probe.

51. (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes;

where the set comprises a first capture probe and a second capture probe;

where the first capture probe and the second capture probe have identical miRNA microRNA binding segment sequences:

where the first capture probe has a first adapter segment sequence that is different from the first adapter segment sequence of the second capture probe; and

where the first capture probe has a second adapter segment sequence that is different from the second adapter segment sequence of the second capture probe.

52. (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes;

where the set comprises a first capture probe and a second capture probe;

where the first capture probe and the second capture probe have identical second adapter segment sequences;

where the first capture probe has a first adapter segment sequence that is different from the first adapter segment sequence of the second capture probe; and

where the first capture probe has an miRNA a microRNA binding segment sequence that is different from the miRNA microRNA binding segment sequence of the second capture probe.

53. (currently amended) The method of claim 33, where the capture probe provided is a set of capture probes;

where the set comprises a first capture probe and a second capture probe;

where the first capture probe has a first adapter segment sequence that is different from the first adapter segment sequence of the second capture probe;

where the first capture probe has an miRNA a microRNA binding segment sequence that is different from the miRNA microRNA binding segment sequence of the second capture probe; and

where the first capture probe has an miRNA <u>a microRNA</u> binding segment sequence that is different from the <u>miRNA microRNA</u> binding segment sequence of the second capture probe.

 (original) The method of claim 33, where the capture probe provided is a set of capture probes;

where the set comprises a first capture probe having a first capture probe sequence, a second capture probe having a second capture probe sequence, and a third capture probe having a third capture probe sequence;

where the first capture probe sequence is different from the second capture probe sequence;

where the first capture probe sequence is different from the third capture probe sequence; and

where second capture probe sequence is different from the third capture probe sequence.

- 55. (original) The method of claim 33, where the first linker segment and the second linker segment comprise a substance selected from the group consisting of one or more than one type of polynucleotide, one or more than one type of polynucleotide analog, and a combination of one or more than one type of polynucleotide analog.
- 56. (original) The method of claim 33, where the first linker, or the second linker, or both the first linker and the second linker are resistant to nuclease degradation.
- 57. (original) The method of claim 56, where the first linker, or the second linker, or both the first linker and the second linker comprise nuclease resistant nucleotides.
- 58. (original) The method of claim 56, where the first linker, or the second linker, or both the first linker and the second linker comprise nucleotides with a phosphothioate backbone

that render the first linker, or the second linker, or both the first linker and the second linker resistant to nuclease degradation.

- 59. (original) The method of claim 56, where the first linker, or the second linker, or both the first linker and the second linker comprise nuclease resistant nucleotides and comprise nucleotides with a phosphothioate backbone that render the first linker, or the second linker, or both the first linker and the second linker resistant to nuclease degradation.
- 60. (original) The method of claim 33, where the first linker and the second linker, each comprises between 6 and 50 residues.
- 61. (original) The method of claim 33, where the first linker comprises at least 10 residues, and at least 10 residues at the 3' end of the first linker are exactly the complement of the corresponding residues at or near the 5' end of the first adapter segment.
- 62. (original) The method of claim 33 where the second linker comprises at least 10 residues, and at least 10 residues at the 5' end of the second linker are exactly the complement of the corresponding residues at or near the 3' end of the second adapter segment.
- 63. (original) The method of claim 33, where the 5' end of the first linker, or the 3' end of the second linker, or both the 5' end of the first linker and the 3' end of the second linker comprise a label.
- 64. (original) The method of claim 33, where the 5' end of first linker comprises one or more than one residue that extends beyond the 3' end of the first adapter segment after the first linker hybridizes to the first adapter segment.
- 65. (original) The method of claim 64, where the one or more than one residue of the 5' end of first linker that extends beyond the 3' end of the first adapter segment functions as a primer binding site.
- 66. (original) The method of claim 33, where the 3' end of second linker comprises one or more than one residue that extends beyond the 5' end of the second adapter segment after the second linker hybridizes to the second adapter segment.

67. (original) The method of claim 66, where the one or more than one residue of the 3' end of second linker that extends beyond the 5' end of the second adapter segment functions as a primer binding site.

- 68. (original) The method of claim 33, where the sample, the capture probe, the first linker and the second linker are combined simultaneously.
- 69. (original) The method of claim 33, further comprising adding one or more than one RNAse inhibitor to the combination of the sample, the capture probe, the first linker and the second linker.
- 70. (original) The method of claim 33, where the first adapter segment comprises a solid phase binding group, or the second adapter segment comprises a solid phase binding group, or both the first adapter segment comprises a solid phase binding group and the second adapter segment comprises a solid phase binding group; and

where the method further comprises binding the capture probe to a solid phase before or after combining the sample, the capture probe, the first linker and the second linker.

- 71. (original) The method of claim 70, where the solid phase is a plurality of paramagnetic particles.
- 72. (currently amended) The method of claim 70, where the capture probe is bound to a solid phase through the first adapter segment or through the second adapter segment or through both the first adapter segment and the second adapter segment; and

where the method further comprises purifying the capture probes with hybridized first linker, miRNA microRNA of interest and second linker-bound to the solid phase by removing non-hybridized first linkers, second linkers and any other substances that are not bound to the solid phase.

- 73. (original) The method of claim 70, where the solid phase is contained in a vessel comprising a surface and a cap, and where purifying comprises applying a magnetic field to attract the solid phase to the surface of the vessel or the cap of the vessel.
- 74. (currently amended) The method of claim 33, where the first linker hybridizes to the first adapter segment at a position where the last residue on the 3' end of the first linker

hybridizes to a residue on the first adapter segment that is between 1 residue and 5 residues from the 3' end of the miRNA microRNA binding segment.

- 75. (currently amended) The method of claim 33, where the first linker hybridizes to the first adapter segment at a position where the last residue on the 3' end of the first linker hybridizes to a residue on the first adapter segment that is immediately adjacent to the 3' end of the miRNA microRNA binding segment.
- 76. (currently amended) The method of claim 33, where the second linker hybridizes to the second adapter segment at a position where the last residue on the 5' end of the second linker hybridizes to a residue on the second adapter segment that is between 1 residue and 5 residues from the 5' end of the miRNA microRNA binding segment.
- 77. (currently amended) The method of claim 33, where the second linker hybridizes to the second adapter segment at a position where the last residue on the 5' end of the second linker hybridizes to a residue on the second adapter segment that is immediately adjacent to the 5' end of the miRNA microRNA binding segment.
- 78. (original) The method of claim 33, where the method further comprises purifying the complex.
- 79. (original) The method of claim 33, where the complex is bound to a solid phase through the first adapter segment or through the second adapter segment or through both the first adapter segment and the second adapter segment; and

where the method further comprises purifying the complex by removing non-hybridized first linkers, second linkers and any other substances that are not bound to the solid phase.

- 80. (currently amended) The method of claim 33, where the method further comprises purifying the ligated first linker-miRNA microRNA of interest-second linker that has been dehybridized from the capture probe.
- 81. (currently amended) The method of claim 80, where the first linker, or the second linker, or both the first linker and the second linker comprise nuclease resistant nucleotides, or comprise nucleotides with a phosphothioate backbone that render the first linker, or the second linker, or both the first linker and the second linker resistant to nuclease degradation; and

where purifying the ligated first linker-miRNA microRNA of interest-second linker comprises applying DNAase to a solution containing the ligated first linker-miRNA microRNA of interest-second linker to destroy any DNA present in the solution.

- 82. (currently amended) The method of claim 80, where purifying the ligated first linker-miRNA microRNA of interest-second linker comprises circularizing the ligated first linker-miRNA microRNA of interest-second linker.
- 83. (currently amended) A method for identifying an miRNA a microRNA of interest, the method comprising:
  - a) isolating the miRNAs microRNAs according to claim 33; and
- b) sequencing the miRNA microRNA of interest portion of the strand of the ligated first linker-miRNA microRNA of interest-second linker.
- 84. (currently amended) The method of claim 83, where sequencing comprises subjecting the strand of the ligated first linker-miRNA microRNA of interest-second linker to reverse transcription to produce a double stranded product comprising a first strand of the ligated first linker-miRNA microRNA of interest-second linker and a second strand that is the complement of the first strand.
- 85. (original) The method of claim 83, where sequencing comprises amplifying the double stranded product to produce amplification products.
- 86. (original) The method of claim 84, where sequencing comprises cloning the amplification products and culturing the amplification products.